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We claim:

- A process for making bioactive glasses comprising:
   preparing a reaction sol capable of forming a sol-gel;
   aging said reaction mixture;
   near equilibrium drying a gel resulting from the reaction mixture, and;
   heating the near equilibrium dried gel at a temperature above ambient.
- 2. The process of Claim 1, further comprising grinding the near equilibrium dried gel.
- 3. The process of Claim 2, further comprising classifying the ground near equilibrium dried gel to various particle size ranges.
- 4. The process of Claim 1, said aging conducted at a temperature of at least about 40°C.
- 5. The process of claim 1, said aging conducted for a duration of at least about 35 hours.
- 6. The process of Claim 1, said near equilibrium drying conducted at about 60 to 98% humidity.
- 7. The process of Claim 1, said near equilibrium drying conducted at a temperature of between about 130°C to about 180°C over at least part of the duration of the near equilibrium drying step.

- 8. The process of Claim 1, said near equilibrium drying conducted at temperatures varied over time between about 130°C to about 180°C.
- 9. The process of Claim 8, said near equilibrium drying conducted at a temperature ramp with a positive time vs. temperature slope over at least part of the duration of said near equilibrium drying step.
- 10. The process of Claim 1, the heating step conducted at between about 200°C to about 700°C over at least part of the duration of the heating step.
- 11. The process of Claim 1, the hearing step conducted at temperatures varying over time between 200°C to about 700°C over at least part of the duration of the heating step.
- 12. The process of Claim 17, the heating step conducted at a temperature ramp with a positive time vs. temperature slope over at least part of the duration of said heating step.
- 13. The process of Claim 1, the reaction mixture including water, hydrochloric acid, tetraethoxysilane, triethylphosphate, or calcium nitrate, or mixtures thereof.
- 14. A near equilibrium dried bioactive glass comprising a silicon dioxide based composition prepared by a sol-gel process capable of forming hydroxycarbonate apatite layer when exposed to physiological fluids.

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The bioactive glass of Claim 14, said glass having an average pore size greater than about 60A° when the silicon dioxide content of the bioactive glass is in the range of about 55 to about 65% by weight.

The bioactive glass of Claim 1, said glass having an average pore size greater than 70A° where the silicon dioxide content is in the range of about 65 to about 75% by weight.

The bioactive glass of Claim 14, said glass having an average pore size greater than 30A° when the silicon dioxide is in the range of about 75 to about 85% by weight.

- 18. A near equilibrium dried bioactive glass composition comprising a silicon dioxide based composition having a pore size greater than a corresponding non-near equilibrium-dried bioactive glass.
- 19. A process for making bioactive glasses by a sol-gel process the improvement comprising near equilibrium drying a sol-gel.
- 20. The process of Claim 19, wherein said near equilibrium drying is accomplished at about 60 to 98% humidity.
- 21. The process of Claim 19 the improvement further comprising near equilibrium drying at a temperature up to about 150°C.

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22. The process of Claim 19, the improvement further comprising near equilibrium drying at an initial temperature of less than about 100°C and a final temperature of less than about 150°C.

23. A near equilibrium-dried bioactive glass comprising, by weight %:

	SiO <sub>2</sub>	-	40 - 90
\	CaO	-	4 - 45
\	\Na₂O	-	0 - 20
	P <sub>2</sub> O <sub>5</sub>	-	2 - 10
	Car	-	0 - 25
	$B_2O_3$	-	0 - 10

and a surface area greater than the non-near equilibrium-dried bioactive glass having an identical composition.

The near equilibrium-dried bioactive glass of Claim 25 wherein said near equilibrium-dried bioactive glass has a surface area of greater than about 175m<sup>2</sup>/g and a silicon dioxide content of about 55 to 65% by weight.

The near equilibrium-dried bioactive glass of Claim 23; wherein said near equilibrium-dried bioactive glass has a surface area of greater than about 250 m<sup>2</sup>/g and a silicon dioxide content of about 65 to 75% by weight.

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- 26. The near equilibrium-dried bioactive glass of Claim 25, wherein said near equilibrium-dried bioactive glass has a surface area of about 300 m<sup>2</sup>/g and a silicon dioxide content of about 75 to 85% by weight.
- 27. A sol-gel process for making a bioactive glass monolith comprising:

preparing a reaction mixture capable of forming a bioactive sol-gel monolith; casting the reaction mixture into a mold of desired shape; aging said reaction mixture cast in said mold at a temperature elevated above ambient; near equilibrium drying the reaction mixture, and;

heating the reaction mixture.

- 28. The process of Claim 27, wherein the reaction mixture comprises deionized water, hydrochloric acid, nitric acid, tetraethoxysilane, triethylphosphate or calcium nitrate or mixtures thereof.
- 29. The process of Claim 27, further comprising conducting a pre-aging step before said aging, wherein said pre-aging comprises aging the reaction mixture at ambient temperature.
- 30. The process of Claim 27, further comprising removing a pore liquor after said aging and before said near equilibrium drying.

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- 31. A near equilibrium-dried sol-gel monolith comprising the product of the process of Claim 27.
- 32. A method for treating orthopedic defects comprising contacting an orthopedic defect with an defect healing amount of pear equilibrium dried sol-gel bioactive glass.
- 33. A composition for the treatment of orthopedic conditions comprising a bioactive sol gel glass capable of forming an HCA layer within 12 hours of exposure to simulated body fluids.
- 34. The composition of claim 33, wherein said glass is capable of forming an HCA layer within 5 hours of exposure to simulated body fluids.
- 35. The composition of claim 33, wherein said glass is capable of forming an HCA layer within 2 hours of exposure to simulated body fluids.
- 36. The composition of claim 33, wherein said glass is more than 50% resorbed 8 weeks after implantation into a patient.
- 37. The composition of claim 33, wherein said glass further comprises at least 77% silicon dioxide.